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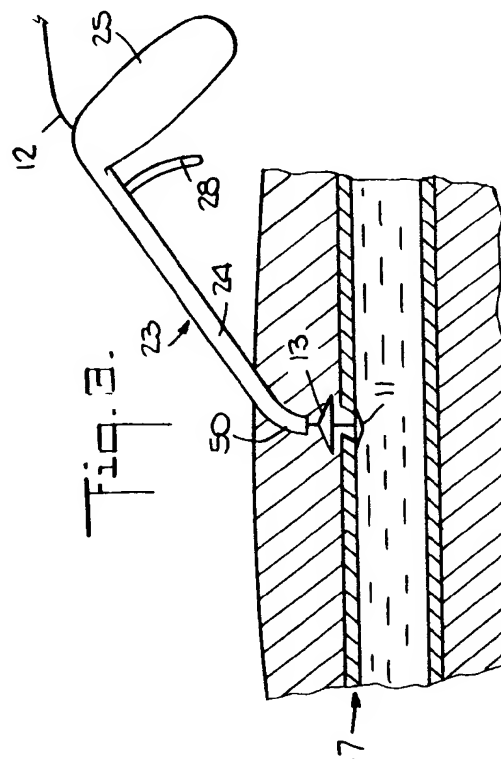
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A sealing device and an insertion tool therefor.

The sealing device is disclosed which is composed of an intra-arterial occluder (11) and an extra-arterial occluder (13), both preferably made of resilient biocompatible and/or bioabsorbable material and held in place via a toothed guide (12) extending integrally from the intra-arterial occluder (11). An insertion tool (23) may also be provided to effect sliding of the extra-arterial occluder (13) over the guide (12) and fixation of the extra-arterial occluder (13) to the guide in sealed relation over an opening in body tissue and safe cutting of the guide. A force gauge (47) is provided on the tool to indicate the sealing pressure on the intra-arterial occluder (11).



This invention relates to a sealing device, for example a percutaneous arterial puncture sealing device, and an insertion tool therefor. More particularly, this invention relates to a sealing device for sealing an opening in a tissue and to a tool for implanting part of the device.

Heretofore, various interventional procedures have been performed which require formation of a puncture or the like in a tissue, for example in a vessel for the introduction of various devices, such as catheters. In this respect, access to arterial and venous vascular systems is necessary for both diagnostic and therapeutic medical procedures. For example, diagnostic arteriography is a radiological procedure which permits visualization of the arterial system for the diagnosis of disease in various organ systems. The most request applications are cardiac catheterization, peripheral vascular angiography, mesenteric angiography, and cerebral angiography. Therapeutic trans-arterial procedures, such as percutaneous transluminal coronary angioplasty (PTCA), have expanded the use of arterial access even further.

All these techniques involve cannulation of an artery so that a catheter may be inserted and advanced into the arterial system. Radio-opaque dye can be injected through the catheter and into the arterial system being examined while X-ray images are recorded. Alternatively, various transcatheter therapies can be employed in the artery, using balloon dilation devices, atherectomy devices, lasers, intra-aortic counterpulsation devices and left ventricular assist devices.

The femoral artery at the junction of the thigh and the abdomen is the most frequent arterial puncture site, but the carotid artery in the neck, the brachial artery in the midarm, and the axillary artery are also used. A percutaneous sheath is usually used whenever multiple catheters are used.

When the sheath is removed, the 1.5 to 5.0 millimetre hole in the artery would spurt blood with significant blood loss unless certain measures are taken. To allow sealing of the hole with a solid clot, heavy pressure is applied to the groin, either manually or with a large C shaped clamp for 10 to 30 minutes. Afterwards, the patient must lie flat on his/her back for 6 to 8 hours, and is usually kept in the hospital overnight for observation. Movement at the hip joint and any increase in intra-abdominal pressure, such as with coughing or bowel movement, can result in serious bleeding. There is a much greater tendency for serious bleeding to occur in patients given anticoagulants like aspirin and heparin or clot-dissolving drugs like streptokinase or tissue plasminogen activator.

Over one million diagnostic and therapeutic arteriographic procedures are done annually in the United States of America. The extra hospital days incurred purely to control bleeding from the arterial puncture site could be avoided if a safe, easy to use and effective arterial puncture sealant device were avail-

able. Such a device would allow same day discharge after angiographic procedures, with a savings of hundreds of millions of dollars. In addition to dollars saved, morbidity from bleeding and complications of blood transfusion could be significantly reduced or virtually eliminated.

Various suggestions have also been made, such as described in U.S. Patents 4,744,364; 4,852,568 and 4,890,612, for the mechanical sealing of such punctures. In such cases, a sealing device in the form of an expandable closure member is to be inserted through a puncture in the vessel, expanded while within the vessel and then retracted against and through the puncture by means of a retraction filament. Thereafter, the filament is left extending from the site of the puncture and through the skin of the patient while being secured in position on the skin of the patient as by a strip of conventional tape. However, such devices permit the exposed thread to be a site for infection. Further, anchorage of the enclosed member in pace by a thread which passes through the skin of a patient may not be reliable so that bleeding may occur by accidental displacement of the sealant device. Further, displacement of the sealant device into the artery would result in arterial occlusion and gangrene.

According to one aspect of the present invention, there is provided a sealing device comprising a first member for positioning on one side of a puncture in a tissue for closing over the puncture, and a guide means extending from the first member for passage through the puncture, the first member being circumferentially collapsible for passage through the puncture and resiliently expandable; characterized in that there is also provided a second member on the guide means for positioning on the opposite side of the puncture to seal the puncture, the second member being resiliently expandable from a circumferentially collapsible state.

Preferably the first member and the second member are made of at least one of a biocompatible material and a bioabsorbable material.

Preferably both the first member and the second member are made of bioabsorbable material and the first member has a faster rate of absorption than the second member.

Preferably at least one of the first and second member includes at least one of an anti-thrombotic material and a clot-promoting material therein.

The guide means is conveniently a guide wire. Preferably the guide wire has a portion containing a plurality of teeth inclined to tent to prevent the second member against movement in a direction away from the first member.

The second member may be of larger outer dimensions than the first member when in the expanded condition.

The present invention makes it possible reliably

to seal a puncture in a vessel or other tissue, in a simple manner.

The first biocompatible member may be a collapsible intro-arterial occluder, for positioning in a vessel to one side of an opening or puncture therein, and the second biocompatible member may be an extra-arterial occluder, which is movably mounted on the guide for positioning outside the vessel on an opposite side of the opening therein and in opposition to the first member in order to seal the opening.

The guide, which may be constructed in the form of a wire, should be made of biocompatible material, and can be integral with the intra-arterial occluder. Further, the guide may be provided with a plurality of teeth near the distal end region so that the extra arterial occluder can pass over the toothed portion in a direction towards the intra-arterial occluder while being retained against movement in the opposite direction.

The construction of the sealing device is such that each of the first and second members is circumferentially collapsible while being resiliently expandable.

The extra-arterial occluder is preferably constructed of larger outer dimensions than the intra-arterial occluder. Further, the intra-arterial and extra-arterial occluders can be made from biocompatible materials which have displayed their safety in humans, including but not limited to Dacron, nylon, Gortex or Teflon. The guide could be made from wire, but other synthetic materials such as a nylon would be applicable or even preferable.

Alternatively, and probably preferably, bioabsorbable materials for the occluders and guide may be ideal, because they would slowly dissolve and leave no residual foreign material behind. Bioabsorbable materials should be chosen so that the rate of absorption of the various parts is differential. Thus, the intra-arterial occluder and guide should be absorbed before the extra-arterial occluder to reduce the chance that the intra-arterial occluder could come free and embolize into the circulation.

Anti-thrombotic materials and/or drugs could be also used or incorporated into the blood contacting surface of the occluders. The occluders and guide may be radio-opaque so that they can be visualized fluoroscopically. Drugs or materials which promote coagulation could be used where the sealing surfaces exist.

The present invention also provides a tool for assisting in the insertion of the sealing device in a patient.

According to a second aspect of the present invention there is provided in combination a sealing device according to the first-mentioned aspect of the present invention and a seal insertion tool for use in placing the second member against the opposite side of the puncture, the seal insertion tool comprising:

a barrel;
a handle fixed to one end of the barrel;
a positioner member slidably mounted in the barrel; and

5 a trigger pivotally mounted with respect to the handle;

wherein the positioner member partially defines a chamber at one end of the barrel to receive the resiliently expandable second member which may serve as an extra-arterial occluder, and the trigger has one end disposed in the barrel; and

10 wherein there is means in the barrel between the one end region of the trigger and the positioner member for pushing the second member in a direction out of the barrel in response to pivoting of the trigger to expel the second member therefrom.

A preferred arrangement is that wherein the positioner member is hollow to define a bore for passage of an elongated guide means which is preferably a guide wire, extending from the first member which may serve as an intra-arterial occluder, through the extra-arterial occluder or other second member; and wherein there is a clamp within the handle for passage of the guide wire therethrough and for selectively clamping the guide wire in response to pivoting of the trigger prior to movement of the positioner member in the barrel.

20 Preferably there is also included a force gauge on the barrel for indicating a degree of pressure exerted via the clamped guide wire on the intra-arterial occluder or other first member at an end of the wire during movement of the positioner member relative thereto.

30 Preferably the tool includes a cutting means for severing the guide wire within the positioner member in response to retraction of the positioner member into the barrel.

35 Preferably the cutting means includes a pair of pincers in the positioner member disposed on opposite sides of the bore, a cam surface on the member and a cam on the barrel for slidably engaging the cam surface to radially deform the positioner member to effect closing of the pincers together to sever the guide therebetween.

40 Preferably the positioner member has a flexible distal portion containing the pincers therein.

Basically, a preferred embodiment of the insertion tool has several functions. First, the tool is to deploy the extra-arterial occluder into the tissues outside the arterial wall. Second, the tool is to provide for an application of a calibrated pressure to the intra-arterial occluder at the contact point of the occluder with the lumen of the vessel. Third, the tool functions so as to cut the guide upon completion of the positioning of the intra-arterial and extra-arterial occluders.

55 In use, the tool is initially provided in a sterile condition with the extra-arterial occluder contained within the chamber at the distal end of the barrel of the

tool. At the conclusion of an arteriographic procedure, a percutaneous sheath would remain in the vessel, e.g. an artery. Thereafter, the intra-arterial occluder would be pushed through the sheath into the arterial lumen, for example, by an obturator whose length is only minimally greater than the sheath. In order to prevent arterial damage, the sheath is pulled back as the obturator is pushed in. As the intra-arterial occluder exits the sheath, the occluder expands due to the inherent natural resiliency, to normal full size. Once the occluder is in the artery, the position of the occluder can be confirmed by noting a free forward movement of the guide and resistance on pulling back. The sheath is then removed from the patient and the guide is pulled back until the intra-arterial occluder is snug against the luminal side of the arterial wall.

Thereafter, the tool is used to position the extra-arterial occluder. To this end, the guide is threaded through the extra-arterial occluder and tool. The tool is then manipulated to position the extra-arterial occluder outside the arterial wall, the appropriate position at which the extra-arterial occluder is to be deployed from the insertion tool into the patient can be determined by the location of the mark on the guide relative to the insertion tool. Alternatively, a stop can be placed on the guide which abuts the tool and prevents the tool tip from being inserted too close to the artery. At this position, the trigger on the tool is squeezed accomplishing several functions. First, the extra-arterial occluder is expelled from with the barrel. At this time, the extra-arterial occluder expands radially under the natural resiliency of the occluder. In this respect, the extra-arterial occluder may have a web-like perimeter from which struts project to positively engage the tissues as the extra-arterial occluder exits from the insertion tool, encouraging the extra-arterial occluder to open as intended.

Second, the two occluders are pressed together with a pressure regulated by the operator and measured by the tool. Further, since the distal end of the tool is flexible, the distal end is able to conform to a position substantially perpendicular to the artery as the occluders are pressed together.

By making the extra-arterial occluder larger than the intra-arterial occluder, several advantages are obtained. First, the larger size ensures that the extra-arterial occluder cannot accidentally be inserted into the artery. Second, the force applied to hold the two occluders together will be applied to a larger surface area, thus reducing the pressure on the tissues. Thus, an excessively high pressure which could result in death of the underlying tissue (necrosis) is avoided.

Once the occluders are properly seated, the cutting means within the tool can be activated to sever the guide just after the point of exit from the extra-arterial occluder. The tool can then be withdrawn leav-

ing the artery positively sealed. In this case, no part of the sealing device remains external to the patient. Instead, the entire sealing device is implanted within the tissue which heals around the device.

For a better understanding of the present invention and to show how the same may be carried into effect, reference will now be made, by way of example, to the accompanying drawings, in which:-

Fig. 1 illustrates a view at the conclusion of an arteriographic procedure with a sheath in place and an intra-arterial occluder, forming part of the device of the present invention, inserted within an arterial vessel;

Fig. 2 illustrates a view similar to Fig. 1 with the intra-arterial occluder of the sealing device in place against the wall of the vessel and with the sheath withdrawn;

Fig. 3 illustrates a view similar to that in Figs. 1 and 2 with an insertion tool in accordance with the present invention in place coaxially surrounding the guide extending from the intra-arterial occluder;

Fig. 4 illustrates a view similar to Fig. 3 of the tool at a time at which the extra-arterial occluder is being secured to the guide extending from the intra-arterial occluder;

Fig. 5 illustrates a cross-sectional view of an arterial vessel with a sealing device according to the present invention in place;

Fig. 6 illustrates on an enlarged scale a part cross-sectional view of the insertion tool shown in Fig. 3 as constructed in accordance with the present invention;

Fig. 7 illustrates a enlarged view of a sealing device in accordance with the present invention in the region of a puncture in an artery;

Fig. 8 illustrates a view of the sealing device of Fig. 7 in fixed position around an opening in a arterial vessel;

Fig. 9 illustrates a plan view of the intra-arterial occluder shown in Fig. 7; and

Fig. 10 illustrates a plan view of the extra-arterial occluder shown in Fig. 7.

Referring now to Fig. 7 and Fig. 8, the sealing device 10 is constructed of a biocompatible and/or bioabsorbable member in the form of an intra-arterial occluder 11, a guide means in the form of an elongated biocompatible and/or bioabsorbable wire 12 integral with the extending centrally from the intra-arterial occluder 11 and a second biocompatible and/or bioabsorbable member in the form of an extra-arterial occluder 13. As indicated, the guide 12 includes a portion extending from the intra-arterial occluder 11 which contains a plurality of saw teeth 14 while the extra-arterial occluder 13 is provided with an opening 15 (see Fig. 10) through which the guide 12 passes. The saw teeth 14 are wider in diameter than the opening 15 in the extra-arterial occluder 13 so that the ex-

tra-arterial occluder 13 can be passed over the teeth 14 in a direction towards the intra-arterial occluder 11. Thus, the extra-arterial occluder 13 can be advanced until a desired amount of pressure is placed on the tissues, such as the wall 16 of an artery 17, while being retained against movement in an opposite direction.

Each occluder 11, 13 is formed of a material and a shape so as to be circumferentially collapsible, for example from the normal position shown in Figs. 9 and 10. Further, each occluder 11, 13 is made to be resiliently expandable from a collapsed state into the normal positions as shown in Figs. 9 and 10.

As shown in Fig. 9, the intra-arterial occluder 11 is of circular disc shape and is provided with a plurality of radial struts 18 integral with flexible sections 19 therebetween.

As shown in Fig. 10, the extra-arterial occluder 13 of umbrella-like construction having a plurality of radially disposed struts 20 with integral web-like sections 21 therebetween. Further, the struts 20 extend radially outwardly of the web portions 21 so as to engage the tissue outside of the vessel 17 (see Fig. 7) to encourage circumferential opening of the occluder 13.

As indicated in Figs. 9 and 10, the extra-arterial 13 is of larger outer dimensions than the intra-arterial occluder 11.

Both occluders 11, 13 are made of a suitable bio-compatible material which may as well be a bioabsorbable material so as to be absorbed over time. In this respect, the rate of absorption of the intra-arterial occluder 11 is faster than that of the extra-arterial occluder 13 and the guide 12 to reduce the risk of the intra-arterial occluder 11 coming free.

Referring to Figs. 7 and 8, the sealing device 10 is constructed such that the intra-arterial occluder 11 can be placed on the inside of the vessel 17 over an opening or puncture 22 therein while the extra-arterial occluder 13 secured over the outside of the vessel 17 on the opposite side of the opening 22. The degree of pressure between the occluders 11, 13 should be sufficient to seal the opening 22 while at the same time not creating undue pressure on the wall 16 of the vessel 17.

Referring to Fig. 6, the insertion tool 23 is constructed so as to facilitate implantation of the sealing device 10. As illustrated, the tool 23 includes a barrel 24, a handle 25 which is fixed to one end of the barrel 24, a positioner member 26 which is slidably mounted in the barrel 24 to define a chamber 27 at one end for receiving the extra-arterial occluder 13 and a trigger 28 which is pivotally mounted on the handle 25 with one end disposed in the barrel 24. As indicated, the trigger 28 is mounted via a pivot pin 29 which is fixed within a yoke 30 integral with the handle 25.

In addition, a stem or extension 31 extends from the positioner 26 into a recess 32 in one end of the trigger 28 and is provided with an enlarged head 33

within the recess 32. As illustrated, the entrance to the recess 32 is formed by a pair of projections 34 which serve to define a slot through which the extension 31 may pass while preventing outward passage of the enlarged head 33. For purposes of assembly, the projections 34 may be sufficiently flexible to permit introduction of the head 33 or may be formed as separate pieces which are removably secured to the trigger 28 in a suitable fashion to permit assembly of the head 33 within the recess 32.

The extension 31 and head 32 serve as a means for pushing the positioner member 26 in a direction out of the barrel 24 in order to expel the extra-arterial occluder 13 therefrom.

A spring wire 35 is also mounted in the trigger 28 to extend through the recess 32 and upwardly, as viewed, through an appropriate slot in the barrel 24 for purposes as described below. The spring wire 35 abuts against the head 33 on the stem 31 so as to transfer a pushing force on the head 33 and thus the positioner member 26 so as to push the member 26 in a direction out of the barrel 24 in order to expel the extra-arterial occluder 13, therefrom.

The positioner member 26 is hollow so as to define a through bore 36 for passage of the guide 12 therethrough. In addition, a clamp 37 is provided within the handle 25 for selectively clamping the guide 12 passing therethrough in response to pivoting of the trigger 28 and prior to movement of the positioner member 26. As indicated, a means is provided between the trigger 28 and the clamp 37 in order to effect a clamping action. This means includes a guide clamp actuator 38 which is pivotally mounted at one end on a pivot pin 39 secured to the trigger 28. In addition, the actuator 38 is slidably mounted within a recess 40 of the handle 25 and has a recess 41 which contains a cam surface 42 which rides on a fixed pin 43 in the handle 25. In addition, the actuator 38 carries one jaw 44 of the clamp 37 in facing relation to a second jaw 45 of the clamp 37 which is integral with the handle 25. As indicated, each jaw 44, 45 has a plurality of serrations 46 to grip the guide 12.

Upon pivoting of the trigger 28 in a clockwise direction, as viewed, the actuator 38 is slid to the right relative to the jaw 44 and raised via the pin 43 thereby bringing the jaws 44, 45 together about the guide 12.

The tool 23 is also provided with a force gauge 47 on the barrel 24 for indicating a degree of pressure exerted on the clamped guide 12 during movement of the positioner member 26 relative to the guide 12. This force gauge 47 is calibrated in pressure units of millimeters of mercury to reflect the force applied to the surface area of the intra-arterial occluder 11 (see Fig. 7). It is expected that the pressure applied by the operator will be between diastolic and just above systolic blood pressure although the optical pressure necessary should be determined from clinical trials.

The force gauge 47 is constructed as an L-shap-

ed element which is mounted on the end of the trigger 28 and which projects through a slot in the barrel 24. In addition, the exposed leg 48 of the gauge 47 carries a printed calibrated scale (not shown) which is coordinated with the spring wire 35 so as to provide a read out of the force applied to the spring wire 35 during pivoting of the trigger 28 in a clockwise Figmanner as viewed. As indicated in 6, a pair of slots 49 may be provided in the barrel 24, one for the spring wire 35 and one for the gauge 47.

As shown in Fig. 6, the distal end of the barrel 24 is provided with a flexible tip 50 while the positioner member 26 is provided with a flexible tip 51. This permits the tool 27 to bend at the end when the extra-arterial occluder 13 is being implanted.

By way of example, the flexible tip 50 of the barrel 24 may be effected by parallel slots which are cut into the barrel 24 from opposite sides without reaching the mid-line of the barrel 24. This allows the barrel to bend in a plane parallel to the handle 25 of the tool 24. In like manner, the flexible tip 51 of the positioner member 26 may be formed with slots of the same nature. In addition, the surface of the barrel 24 and the positioner member 26 can be lined with a flexible and deformable material which will allow them to slide past each other without undue resistance.

In addition, a cutting means 52 is provided for severing the guide 12 within the positioner member 26 in response to retraction of the member 26 into the barrel 24 after implanting of the extra-arterial occluder 13. As indicated, this cutting means 52 includes a pair of pincers 53 in the flexible positioner tip 51 on opposite sides of the bore 36, a cam surface 54 within a groove 55 of the positioner 26 and a cam 56 fixedly mounted on the inside wall of the barrel 24 for slidably engaging the cam surface 54 in order to radially deform the flexible positioner tip 51 so as to effect closing of the pincers 53 together with sever the guide 12 therebetween.

Referring to Fig. 1, in order to seal the opening 22 in the artery 17, for example at the end of an arteriographic procedure, a percutaneous sheath 57 which is normally used in that procedure is retained in place. The intra-arterial occluder 11 which when deployed is larger than the anticipated opening 22, together with the guide 12 is pushed through the sheath 57 into the lumen of the artery 17 by an obturator (not shown). As the occluder 11 exits the sheath 57, the occluder 11 resiliently expands to full size.

Next, referring to Fig. 2, the intra-arterial occluder 11 is pulled back against the inside wall of the artery 17 and the sheath 57 removed.

Next, the insertion tool 23 is slid over the guide 12. At this time, the collapsed extra-arterial occluder 13 nests in the chamber 27 (see Fig. 6) of the tool 23 while the guide 12 passes through the extra-arterial occluder 13 and through the entire tool 23.

When the trigger 28 on the tool 23 is activated,

two sequential functions occur, one after the other. First, the guide 12 is clamped in place relative to the tool 23 via the clamp 37. In this respect, during initial travel, the trigger 28 moves the clamp actuator 38 forwardly, i.e., to the right as shown in Fig. 6. The actuator 38 then translates upwardly to force the clamp jaws 44, 45 (or other high friction mechanism) together in order to fix the guide 12 in place. During this part of the trigger movement, there is no movement of the positioner member 26.

Next, as the trigger 28 continues to pivot, the end of the trigger 28 pushes the spring wire 35 in contact with the enlarged head 33 and, thus, the positioner member 26. The spring wire 35 then deforms toward the rear of the tool in proportion to the force applied as the positioner member 26 moves forwardly to expel the extra-arterial occluder 13 (Fig. 3). The force which is applied to the positioner member 26 is resisted by the guide 12 which is fixed in place by the guide clamp 37. The resisting force is transmitted through the guide 12 to the intra-arterial occluder 11 which is being pressed against the inner surface of the arterial wall. The force applied divided by the known surface area of the intra-arterial occluder 11 yields the pressure exerted against the wall of the artery by the intra-arterial occluder. This pressure is registered on the calibrated scale printed on the force/pressure gauge 47.

As the travel of the trigger 28 continues, the extra-arterial occluder 13 is advanced along the guide 12 until exiting the tool 23. Thereafter, an increase in spreading force is applied between the extra-arterial occluder 13 and the fixed guide 12 (see Fig. 4). The force on the guide 12 and the positioner member 26 which is registered on the force gauge 47 is effectively transmitted to the intra-arterial occluder 11. As pressure is applied, the flexible tip 50 of the barrel 24 and the flexible tip 51 of the positioner member 26 conform to a shape perpendicular to the wall 16 of the vessel 17 (see Fig. 4).

Once the extra-arterial occluder 13 is positioned, the guide 12 is severed. To this end, the trigger 28 is pivoted in a reverse direction, i.e. counterclockwise as viewed in Fig. 6, so as to retract the positioner member 26 into the barrel 24. To this end, the trigger 28 is provided with a cut-out 57 to accommodate the fingers so that the trigger 28 is easily pulled away from the handle 25.

As the positioner member 26 retracts, the cam 56 engages the cam surface 54 of the flexible positioner tip 51 causing the pincers 53 to come together. The pincers 53 then exert a sufficient force to sever the guide 12 thereat. Since the cutting operation occurs within the positioner member 26, there is little danger of cutting the guide 12 too short which could inadvertently allow the intra-arterial and extra-arterial occluders 11, 13 to separate.

The insertion tool 23 is then removed from the

patient and a band-air or similar dressing is applied to the wound.

Of note, the occluders 11, 13 may be held together at the desired pressure other than through the use of the guide saw teeth 46. For example, glues may be used.

The invention thus provides a sealing device which is fixedly secured in place completely internal in the patient. There are no external members which may leave a path for infection into the patient. Further, sealing device may be used to close percutaneous punctures in any number of body parts including, but not limited to, the gall bladder, stomach, intestine, lung, heart, urinary bladder, urinary collecting systems veins.

Further, the invention provides a tool which can be readily manipulated to implant a sealing device totally within a patient.

Further, the invention provides a relatively simple tool for the deployment of a sealing device about an opening in body tissues.

Still further, the invention provides a tool which is capable of measuring the occluder pressure during insertion.

Claims

1. A sealing device comprising a first member (11) for positioning on one side of a puncture in a tissue for closing over the puncture, and a guide means (12) extending from the first member (11) for passage through the puncture, the first member (11) being circumferentially collapsible for passage through the puncture and resiliently expandable; characterized in that there is also provided a second member (13) on the guide means (12) for positioning on the opposite side of the puncture to seal the puncture, the second member (13) being resiliently expandable from a circumferentially collapsible state.
2. A sealing device as claimed in claim 1, wherein the first member (11) and the second member (13) are made of at least one of a bio-compatible material and a bioabsorbable material.
3. A sealing device as claimed in claim 1, wherein both the first member (11) and the second member (13) are made of bioabsorbable material and the first member (11) has a faster rate of absorption than said second member (13).
4. A sealing device as claimed in claim 1, 2 or 3, wherein at least one of the first and second members (11, 13) includes at least one of an anti-thrombotic material and a clot-promoting materi-

al therein.

5. A sealing device as claimed in any preceding claim, wherein the guide means (12) is a guide wire.
6. A sealing device as claimed in claim 5, wherein the guide wire (12) has a portion containing a plurality of teeth (14) inclined to tend to prevent the second member (13) against movement in a direction away from the first member (11).
7. A sealing device as claimed in any preceding claim, wherein the second member (13) is of larger outer dimensions than the first member when in the expanded condition.
8. In combination, a sealing device as claimed in any preceding claim, and a seal insertion tool for use in placing the second member (13) against the opposite side of the puncture, the seal insertion tool comprising:
 - a barrel (24);
 - a handle (25) fixed to one end of the barrel (24);
 - a positioner member (26) slidably mounted in the barrel (24); and
 - a trigger (28) pivotally mounted with respect to the handle (25);
 wherein the positioner member (26) partially defines a chamber (27) at one end of the barrel (24) to receive the resiliently expandable second member (13) which may serve as an extra-arterial occluder, and the trigger (28) has one end disposed in the barrel (24); and
 - wherein there are means (31, 32) in the barrel (24) between the one end region of the trigger (28) and the positioner member (26) for pushing the second member (13) in a direction out of the barrel (24) in response to pivoting of the trigger (28) to expel the second member (13) therefrom.
9. A combination as claimed in claim 8, wherein the positioner member (26) is hollow to define a bore (36) for passage of an elongated guide means (12) which is preferably a guide wire, extending from the first member (11) which may serve as an intra-arterial occluder (13), through the extra-arterial occluder or other second member (13); and wherein there is a clamp (37) within the handle (25) for passage of the guide wire therethrough and for selectively clamping the guide wire (12) in response to pivoting of the trigger (28) prior to movement of the positioner member (26) in the barrel (24).
10. A combination as claimed in claim 9, which fur-

ther includes a force gauge (47) on the barrel (24) for indicating a degree of pressure exerted via the clamped guide wire (12) on the intra-arterial occluder or other first member (11) at an end of the wire during movement of the positioner member (26) relative thereto. 5

11. A combination as claimed in claim 9 or 10, which further includes a cutting means (52) for severing the guide wire (12) within the positioner member (26) in response to retraction of the positioner member (26) into the barrel (24). 10

12. A combination as claimed in claim 11, wherein the cutting means (52) includes a pair of pincers (53) in the positioner member (26) disposed on opposite sides of the bore (36), a cam surface (54) on the member (26) and a cam (56) on the barrel (24) for slidably engaging the cam surface (54) to radially deform the positioner member (26) to effect closing of the pincers (53) together to sever the guide therebetween. 15 20

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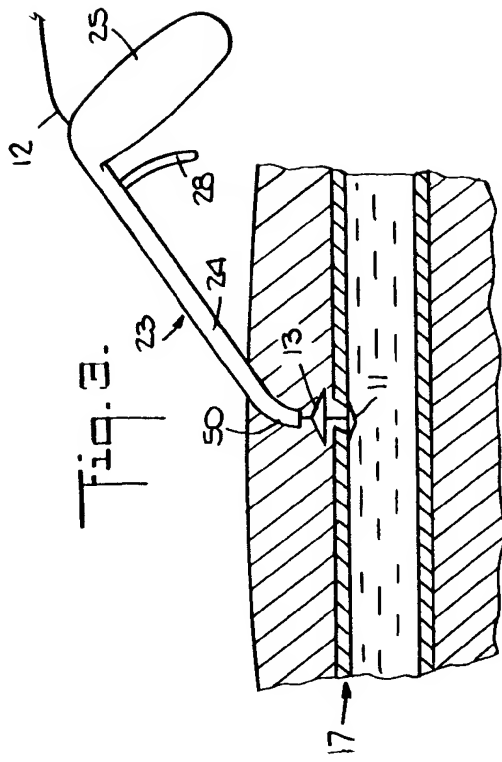
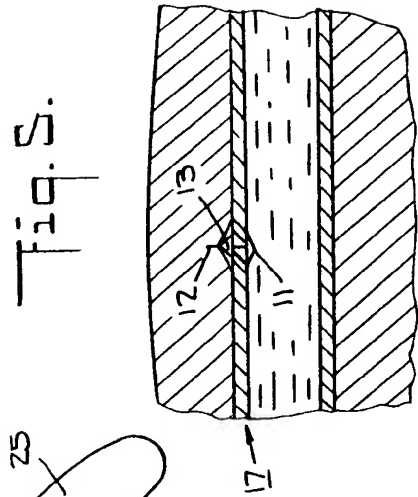
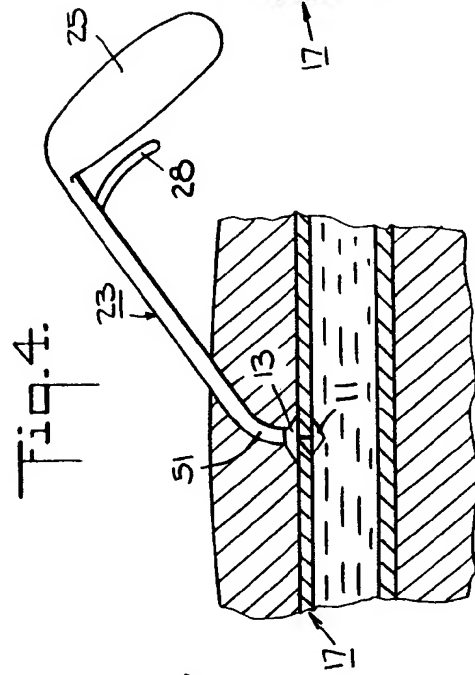
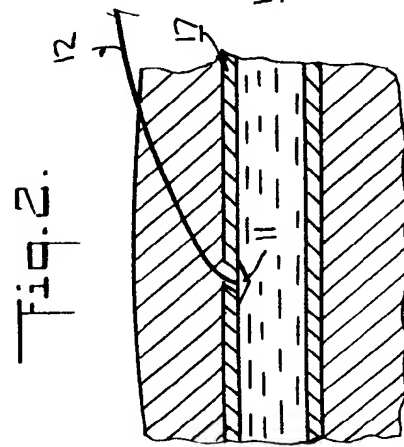
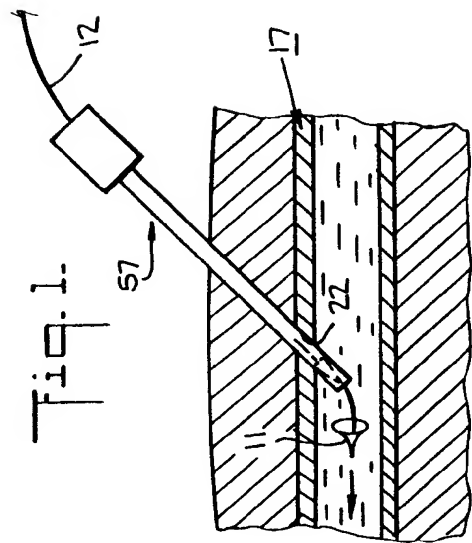
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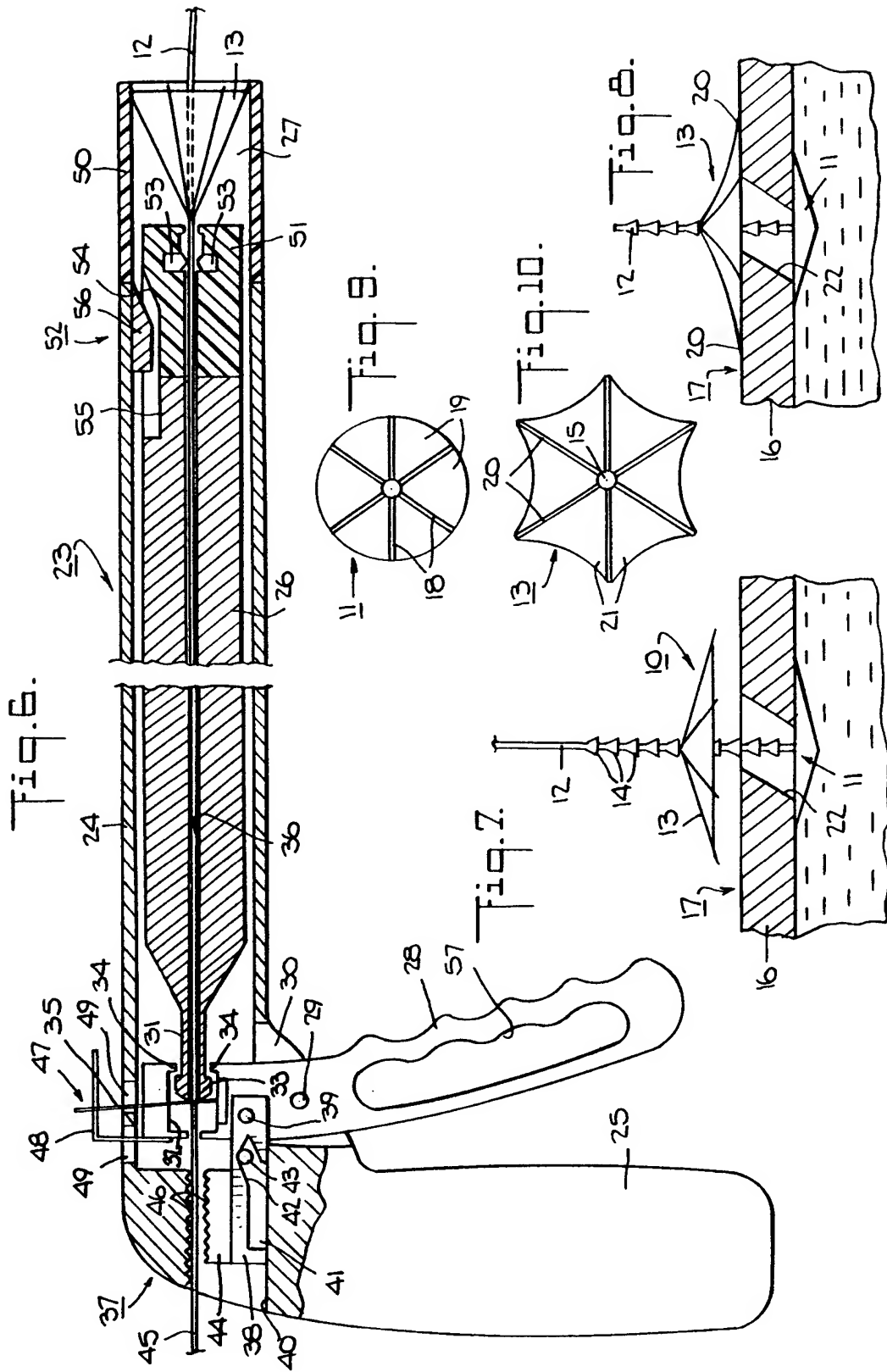
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European Patent
Office

EUROPEAN SEARCH REPORT

Application Number

EP 92 30 8566

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
X	US-A-3 874 388 (KING ET AL.) * column 6, line 8 - line 60 * * column 9, line 9 - column 10, line 16; figures 1A-C, 9A-K * ---	1,2,4,5, 8,10	A61B17/00 A61B17/12
X	WO-A-9 014 796 (MUIJS VAN DE MOER) * page 4, line 12 - page 5, line 15; figures 3,4,6 * ---	1,2,4,5, 6	
A	WO-A-9 116 858 (KENSEY NASH) * page 10, line 3 - line 32 * * page 15, paragraph 2 - page 17, paragraph 1 * * page 18, paragraph 2; figures 1,2,4-7 * & US-A-5 021 059 4 June 1991 ---	8-10	
A	EP-A-0 362 113 (SIDERIS) * column 5, line 4 - line 21; figure 8 * ---	11	TECHNICAL FIELDS SEARCHED (Int. Cl.5)
A	US-A-4 078 305 (AKIYAMA) * abstract; figures 4-6 * ---	11	A61B A61F
A	JP-A-54 009 482 (...) * figure 1 * -----	6	
The present search report has been drawn up for all claims			
Place of search THE HAGUE		Date of completion of the search 22 DECEMBER 1992	Examiner MOERS R.
<p>CATEGORY OF CITED DOCUMENTS</p> <p>X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document</p> <p>T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons & : member of the same patent family, corresponding document</p>			

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